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# Multivalent Cation Crosslinking Suppresses Highly Energetic Graphene Oxide’s Flammability

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## Abstract

Graphene oxide (GO), a common intermediate for making graphene-like materials from graphite, was recently found to possess an explosive fire-hazard that can jeopardize the GO’s large-scale production and wide applications. This work reports a simple and facile method to cross-link the GO with Al<sup>3+</sup> cations, in one step, into a freestanding flexible membrane. This inorganic membrane resists in-air burning on an open-flame, at which non-cross-linked GO was burnt out within ~5 seconds. All characterization data suggested that the in-situ “epoxy ring opening” reactions on GO surface facilitated the cross-linking, which elucidated a new mechanism for the generalized inorganic polymerization. With the much improved thermal- and water-stabilities, the cross-linked GO-film can help to advance high-temperature fuel-cells, electronic packaging, etc. as one of the long-sought inorganic polymers known to date.

## INTRODUCTION

Recently, graphene-based new materials have attracted enormous excitement, thanks to their excellent mechanical and electrical properties on top of their highly accessible specific surface area. In search for better and cheaper routes to synthesizing these materials,<sup>1-7</sup> chemical modification of graphene oxide (GO) is considered the most easily scalable to date<sup>8-16</sup>. This is because, sterically, the modification can easily take place on the oxygenated functional groups (e.g. >O, -OH, -COOH, etc.) that form mostly on the GO edge for exfoliating graphite-layers and dispersing GOs in aqueous and organic solvents. On the GO-surface, the energetic epoxide group was recently found to make the GO highly flammable,<sup>17</sup> and inorganic by-products including potassium and sodium salts (i.e. the residue from the GO synthesis) were shown to contribute significantly to the violent combustability of the GO in ambient air. This fire-hazard makes the GO to be a dangerous material,<sup>18-21</sup> especially for the partially reduced GO (or rGO). Hence, a new method should be developed timely for facilely mass-producing flame-retardant and highly thermal-stable GO.<sup>22</sup>

Herein, we report a new and simple method for mass-producing such non-flammable GO, by cross-linking the GO with Al<sup>3+</sup> cations in one-step in aqueous solutions at room-temperature. The cross-linked GO (cl-GO) resists combustion in ambient air on open-flame, and shows in addition a greatly improved thermal stability, which ended the abovementioned fire-hazard. This thermally stable cl-GO can be applicable to making devices operational at elevated temperatures (above 120 °C) even in air, such as high-temperature fuel cells, high-temperature coating, thermally stable electronic packaging, to name a few. Our characterization data further suggest that the cross-linked GO inherited all characteristics of ordinary GO except the flammability, and its good dispersibility in water being fine-tunable widely, which greatly expands the new cl-GO's processibility and in turn its wide applicability in industry scale.

**EXPERIMENTAL METHODS**

**Chemical synthesis of GO.** The GO was prepared by mixing 0.5 g graphite powder (Alfa Aesar, natural, briquetting grade,-200 mesh,99.9995% metal basis) and 0.5 g NaNO<sub>3</sub> (Alfa Aesar, 98+%) into 23 mL of concentrated H<sub>2</sub>SO<sub>4</sub> (BDH Aristar, 95 –98 % min) solution, under stirring in an ice bath for 15 minutes. This was followed by adding 4 g of KMnO<sub>4</sub> (J.T.Baker, 99% min) gradually under stirring for another 30 minutes in an ice bath, and then transferred into a 40°C water-bath under a stirring for about 90 minutes. The resultant paste was diluted by 50 mL deionized water, then stirred for 15 minutes, and then mixed with 6 ml of H<sub>2</sub>O<sub>2</sub> (Alfa Aesar 29-32% w/w) and 50 mL distilled/deionized (DDI) water. The resultant product was washed with a copious amount of DDI water and dried at 40°C in air over 24 hours.

**Synthesis of aluminum cross-linked graphene oxide (cl-GO).** 300 mg of GO will be dispersed in 100 mL of DDI water under agitation. Separately, 0.2 g of Al(NO<sub>3</sub>)<sub>3</sub>•9H<sub>2</sub>O (EM Science) was added to another 100 mL flask pre-filled with DDI water. The GO dispersion was gradually added into the aluminum nitrate solution, and the resultant cl-GO was stirred for 5 minutes at room temperature, then washed with copious amount of DDI water for several times.

**Fabrication of GO and cl-GO films.** Same amounts of dispersed cl-GO and GO (1 mg/mL) were used to fabricate films on various substrates such as silicon wafer, polystyrene, polyethylene terephthalate, polytetrafluoroethylene, glass slide, and plastic paraffin film. The best defect-free and durable freestanding GO and Al/GO films were formed on polystyrene substrates using drop-casting methods. After the drop casting on the substrates, films were formed from air-drying over 24 hours at room temperature.

**Characterizations.** The GO and cl-GO samples were analysed by means of PHI Versa Probe Scanning X-ray photoelectron spectroscopy (XPS) Microprobe equipped with dual beam charge neutralization and a monochromatic Al K alpha source (1486 eV). Surveys were obtained with an 117eV pass energy and 1.0 eV step size, while high resolution spectra were obtained with 23.5 eV pass energy with 0.1-0.2 eV step size, and with the time of 25 ms per step. For charge correction, 284.8eV was used as the adventitious carbon peak position, and peak positions were determined by the curve-fitting method.

Thermogravimetric Analysis (TGA) tests were performed on TGA Q50 V20.10 Build 36 under N<sub>2</sub> flow, after the samples being heated from room-temperature to 350 °C at the ramping speed of 15 °C/min. The Differential Scanning Calorimetry (DSC) results were obtained in a N<sub>2</sub>-flow (20 ml/min) on Perkin Elmer Pyris Diamond Differential Scanning Calorimeter for 5 mg of each sample, first being heated at 50°C for 1 minute then heated up to 300°C at a speed of 10°C /min. The X-ray Powder Diffraction (XRD) patterns were obtained from a Rigaku MiniFlex II Desktop XRD using monochromatized Cu-K $\alpha$  radiation ( $\lambda = 1.5418$  Å) at 30 kV and 15 mA, in the range of 2-theta from 5° to 60° at a speed of 0.1°/min. High Resolution Scanning Electron Microscopy (SEM) images were obtained using a FEI Nova Nanolab 200 Duo-Beam Workstation being operated on a 15 kV electron beam. In-house built Raman spectroscopy equipped with 532nm laser source at 3mW was used to obtain the microRaman spectra. For estimating surface area using the Methylene Blue Absorption Method, the known masses of GO and cl-GO were separately soaked into an aqueous solution of metylene blue in 25 ml flasks, then stirred at 400 rpm for 48 hours, then the samples were centrifuged, and the supernatant's concentration were analysed using Ultraviolet Visible Spectroscopy (wavelength of 661 nm, U-0080D) for comparison against the original concentration and knowing the MB molecules being adsorbed. Transmission electron microscopy (TEM) images were obtained on a JOEL-2011 electron microscope operating at

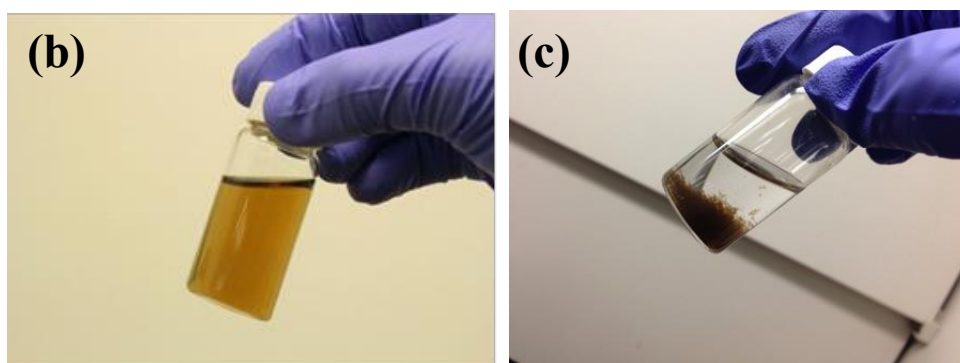
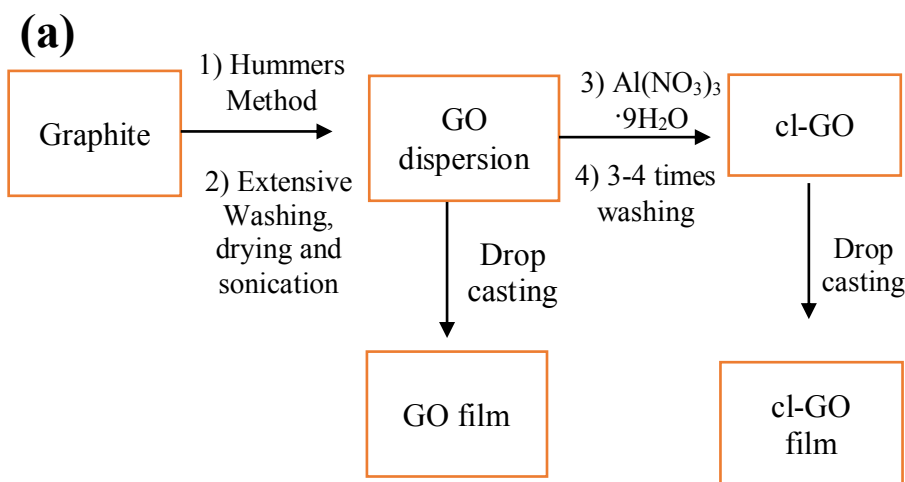
200 kV equipped with an Oxford Link ISIS system for energy-dispersive X-ray spectroscopy (EDX).

## RESULTS/DISCUSSION

It was detailed in literature that even an air-drying temperature near 100 °C can trigger a thermal reduction-decomposition of GO, which is potentially dangerous in large-scale manufacturing.<sup>17</sup> This is because such a decomposition of GO is as highly exothermic as almost self-igniting that must be absolutely avoided in any large-scale manufacturing. Moreover, it was indicated in literature that residual potassium salts, from the GO-synthesis involving KMnO<sub>4</sub> or K<sub>2</sub>S<sub>2</sub>O<sub>8</sub>, can readily transform to various potassium-containing impurities<sup>18</sup> that can help turn the GO to the extremely flammable forms. Removing these impurities by employing filtration or dialysis is time-consuming and costly, because GO-flakes easily clogged the filter-pores and reduced the water-flow across the filtering media (e.g. anodized aluminum oxide or AAO). Washing with abundant water was proven troublesome in our experiment, because after a few washing-cycles the GO started to irreversibly gelatinize which drastically increased the time and manpower in the follow-up centrifugal separations. Further, both the filtration and washing still resulted in the GO-flakes with energetic epoxide groups that can make the GO to be flammable.<sup>17</sup> Thus, an increasing concern has been seriously raised on the fire-hazard of the GO in especially its large-scale production and applications.<sup>19–21</sup>

Our experiment started from the GO-synthesis (Fig 1a) using a modified Hummer's method,<sup>22</sup> and the resultant GO was washed with water and centrifugation. The GO material was then dried in an oven, and thereafter exfoliated in DDI water using an ultra-sonication. The suspension of the exfoliated GO was added into an aqueous solution (1.0% w/w) of Al(NO<sub>3</sub>)<sub>3</sub> under a vigorous stirring, in order for the cross-linking to take place instantly at the room-temperature (Figure 1b-c). This cross-linking was followed by a few times of washing with DDI water, for further reducing the K-containing impurities' content. Afterwards, 100ml of the

GO suspension was centrifuged at 4000 rpm for an hour, and the resultant GO precipitate was collected and then drop-cast on a glass-slide surface to dry into a thin flexible freestanding membrane about 15-20 microns thick. The cl-GO membrane, together with another similar-sized GO-film but without the cross-linking, were each exposed to an open-flame from a commercial lighter (burning the butane-fuel) in air.



**Figure 1.** a) The flowchart for fabricating the GO and cl-GO; b) A GO solution (0.5mg/ml); c) 1 minute after the GO being cross-linked in the aqueous solution (1.0 wt%) of  $\text{Al}(\text{NO}_3)_3$ .

In chemical science, alkaline earth metal cation is a fairly strong Lewis acid that can form a strong bond on GO, by inducing a ring-opening reaction<sup>1</sup> of the epoxide (a Lewis base) on the GO. The epoxide groups are mainly accountable for the energetic behavior of GO, hence the ring-opening reaction on epoxide group can alter the thermal decomposition kinetics.<sup>17</sup> This motivated us to propose logically and prove in experiment whether this concept's applicability

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2  
3 138 could be expanded to using trivalent metal cations such as  $\text{Al}^{3+}$  (a much stronger Lewis acid)  
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5 139 to bond with oxygen-containing functional groups including epoxide and carboxylic acid  
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8 140 groups in between two adjacent GO-sheets, in either a face-to-face or a shoulder-by-shoulder  
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10 141 manner or even both.  
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12  
13 142 Combustion rapidly propagating made the GO-film to vanish (or gasify) in ~5 seconds,  
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15 143 while no combustion (besides reduction) took place on the cl-GO film even after a minute (see  
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17 144 videos in the Supporting Information). Since the open flame is an easily accessible heat source  
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19 145 for GO reduction in scale-up production, we were motivated to further investigate the thermal  
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21 146 behavior for the GO-film and the cl-GO-film.  
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24 147 Thermogravimetric Analysis (TGA) data of GO and cl-GO were compared in Figure  
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26 148 2a. A minor mass-loss for both samples at 100 °C can be attributed to the desorption of  
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28 149 physisorbed water on the samples, while the major mass-losses at 100 °C–300 °C are due to  
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30 150 the pyrolysis of the oxygen-containing functional groups. The cl-GO exhibited a slower mass-  
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32 151 loss starting around 200 °C, while that of the GO appeared at 125 °C in a faster rate.  
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36 152 Differential scanning calorimetry (DSC) results (Figure 2b) further suggested that the  
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38 153 GO's thermal decomposition process is much more exothermic than the cl-GO's. Intuitively,  
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40 154 the excessive and abrupt heat-release of GO from the deoxygenation reaction can trigger the  
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42 155 combustion. In contrast, the heat effect of the cl-GO was much smaller. The DSC data, together  
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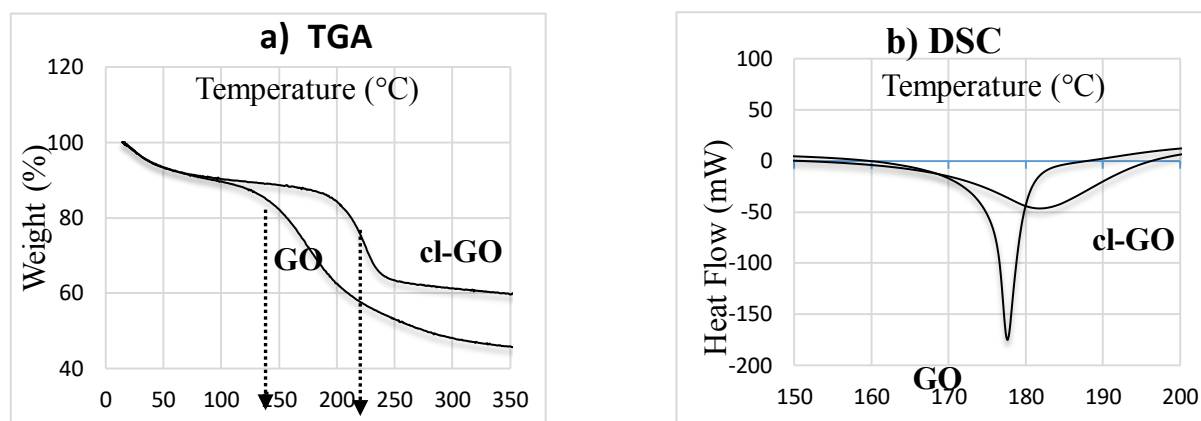
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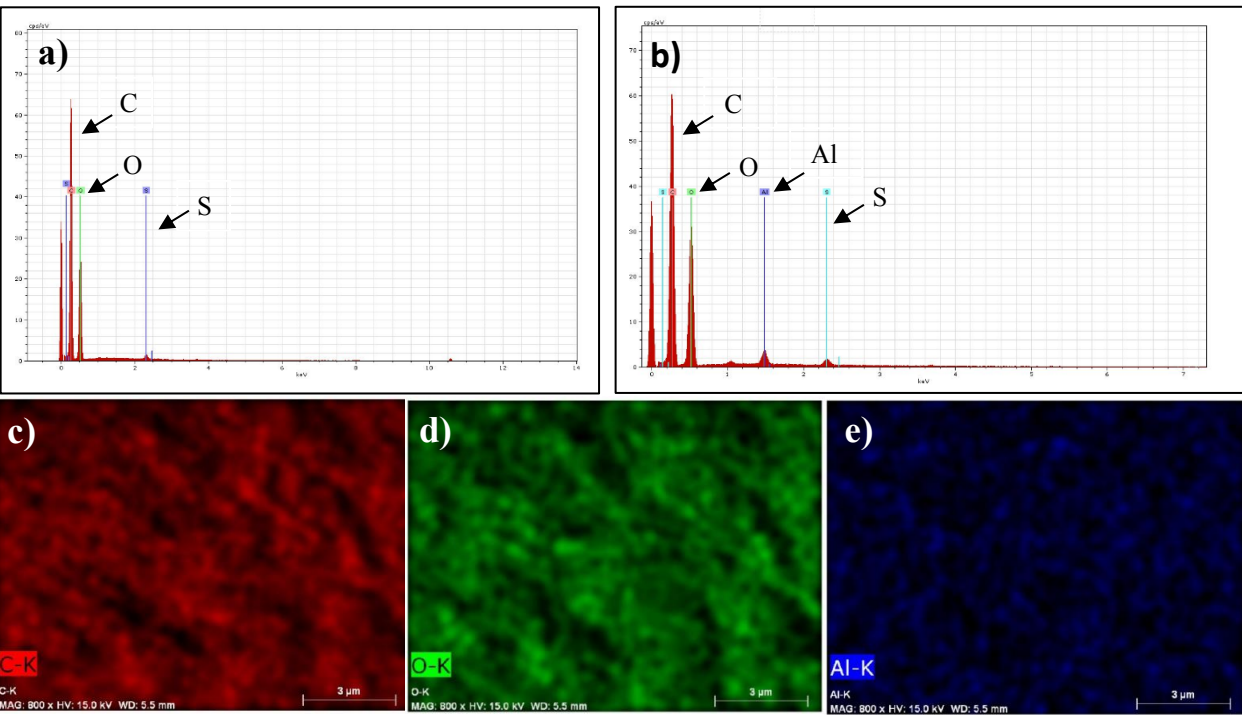
**Figure 2.** a) TGA curves of GO and cl-GO, from a heating at 15 °C/min under a N<sub>2</sub>-flow, showing the mass-loss for cl-GO near 200 °C and that for GO near 125°C. b) DSC curves of GO and cl-GO, from a heating at 10 °C/min under a N<sub>2</sub>-flow, showing the heat release (the exothermic peak) of the energetic GO being much greater than that of the thermal-stable cl-GO.

In thermochemistry, GO's decomposition can shift to the more exothermic site due to an increased content of epoxide, and to the more endothermic due to an increased hydroxyl content.<sup>17</sup> The TGA-DSC data suggest that cross-linking Al<sup>3+</sup> cations on every GO sheets triggered the epoxide ring-opening reaction which decreased epoxide group's content and in turn increased the hydroxyl group content on cl-GO.

For further verifying potassium and sulfur salts' role in the flammability of GO,<sup>21</sup> a GO-film and a cl-GO film were soaked into a 1.0 wt% aqueous solution of KOH for 5 minutes, and then dried and open-flamed. Again, the GO-film was ignited instantly and disappeared quickly, while cl-GO film was not combusted but turned into a reduced cl-GO. This study further concluded the cl-GO film's nonflammable nature.

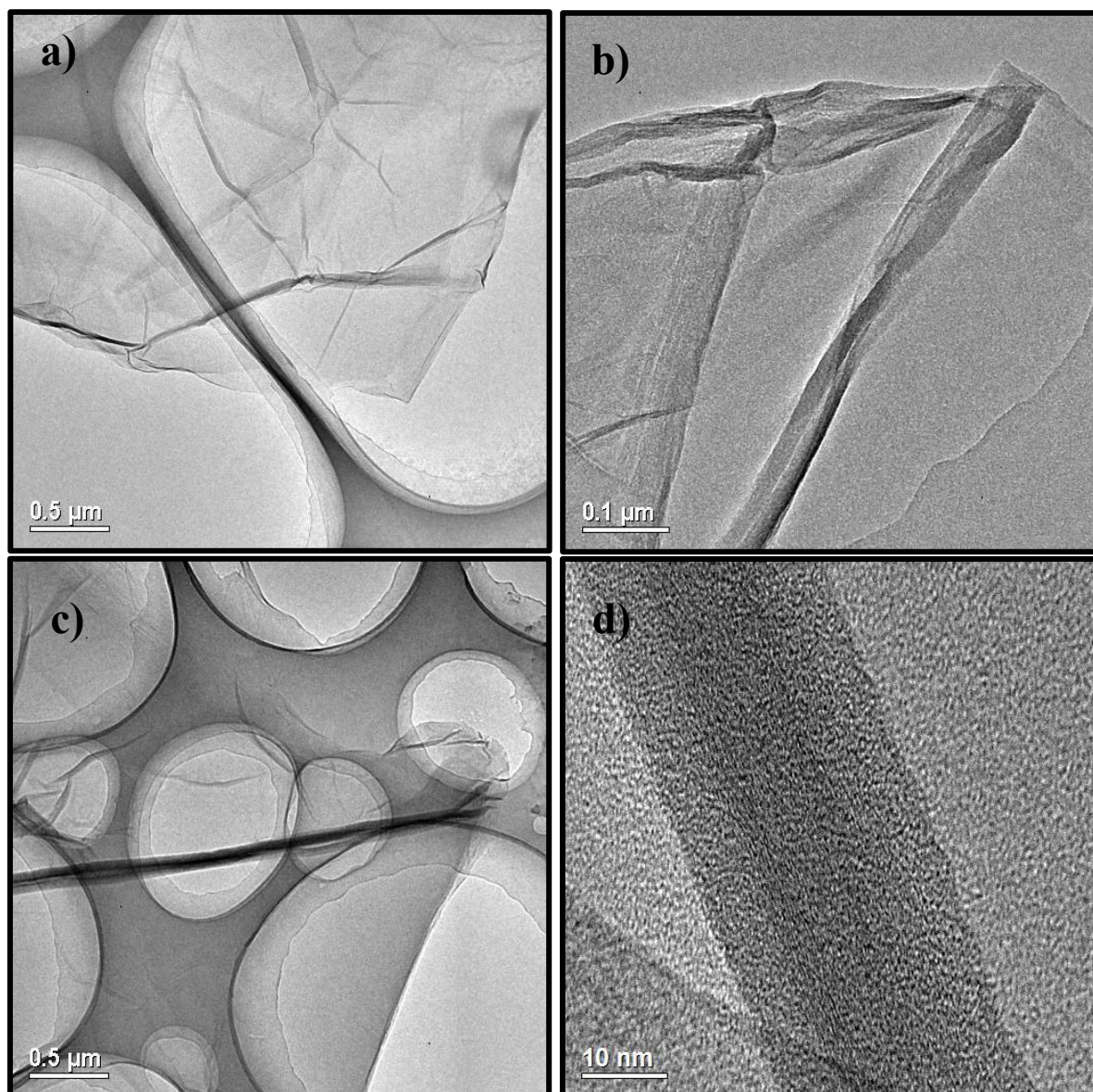
An elemental analysis revealed that GO and cl-GO films have sulfur-content of 0.17 at.% and 0.21 at.%, respectively even after being washed extensively with DI water under centrifugation (Figure 3). Surprisingly, as little as 0.42 at% aluminum has led to the GO-polymerization into the cl-GO within a few seconds. Elemental analysis mapping results of cl-

GO also showed that Al sparsely distributed along the flakes.



**Figure 3.** Elemental analysis results from a) GO; and b) cl-GO; and the supporting elemental mapping of c) carbon; d) oxygen; e) aluminum.

The transmission electron microscopic (TEM) image in Figure 4a disclosed the wrinkled nature of graphene sheets. Figure 4b-c further show that the cl-GO sheets are linked up on the GO-edges, and the split ends of two adjacent sheets can be seen in Figure 4c. Under a higher magnification (Figure 4d), a darker middle section is probably due to the existence of a higher content of Al elements.



**Figure 4.** TEM image of cl-GO; a) Characteristic wrinkled GO-sheet; b) and c) Few layers of overlapped and cross-linked GO-sheets; d) High resolution image of the cl-GO sheets.

New reports in literature have indicated that epoxide groups of GO are mainly responsible for GO's high flammability and alteration of these groups can minimize the energetics<sup>17,18,21</sup>. Thus, we proposed that reducing epoxide groups or all oxygenated functional groups of GO may further minimize the fire hazard. However, it was argued that even the reduced GO (called rGO) can still be extremely flammable due to the residue by-products from the synthesis process of GO<sup>18</sup>. On the other hand, the applications can be limited if all

oxygenated functional groups are removed in GO. Thus, cl-GO possess unique advantages, by eliminating high flammability of GO without disturbing major portion of oxygenated groups. The flame resistance and retardancy of cl-GO can be explained by the two major mechanisms: i) partial pacification of epoxide groups via cross linking them with  $Al^{+3}$  cations, ii) shielding heat propagation between the GO flakes so to result in the cl-GO with superior thermal stability and open flame resistance. One such study reported that upon heating GO the heat transferred from one highly energetic GO' site to another very rapidly<sup>18</sup>, which matches our results from this work. Our DSC results suggested that when once GO was heated, a thermal decomposition started to propagate quickly with rapid heat accumulation which is in line with others reports on the same phenomena. However, when cl-GO was heated, much less heat accumulation between GO flakes prevented the excessive heat accumulation that results in serious fire-hazard.

Surprisingly, some studies in the literature claimed that GO and its derivatives can be used as a potential flame retardant polymer additive despite its highly energetic structure and thermal instability<sup>23–29</sup>. In contrary, a recent review on GO's thermal instability seriously questioned its use for flame retardant applications and pointed out that GO or contaminated rGO may behave like a fuel for combustion rather than a flame retardant additive material<sup>21</sup>. The results in the review suggest that there is a strong association between the number of epoxide groups, the amount of synthesis by-products and GO's flammability. For timely clarifying the confusion in literature, our cl-GO can be a very strong candidate to resolve the abovementioned issues with its thermal stability and flame resistant properties.

Freestanding flexible films made of GO were demonstrated to be highly promising in many important applications in solutions,<sup>30–35</sup> owing to their unique properties such as high tensile strength, proton conductivity, and durability in water. In literature,<sup>36–40</sup> however, negatively charged GO was found soluble in water owing to the presence of the residual metal cations, and  $Al^{3+}$  on the anodized aluminum oxide filter was reported to improve the GO sheets

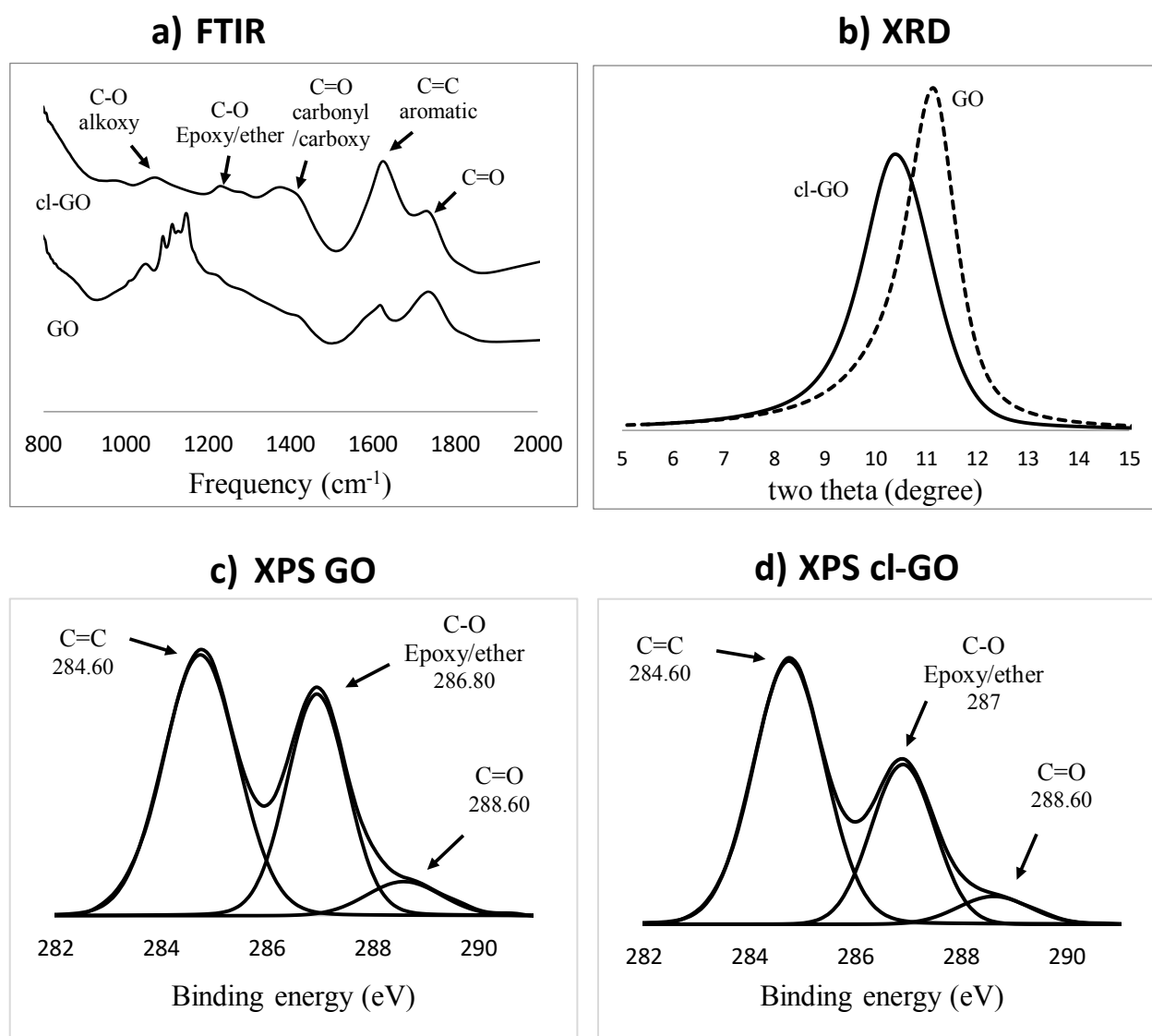
mechanical strength.<sup>40</sup> Hence, in our experiment by sonicating a GO film and a cl-GO film in water for 10 minutes, the GO film became fragmented while the cl-GO film remained intact, which defined a need to study the cl-Go's unusual microstructure.

The cross-linking was further supported by the Fourier-transform infrared (FT-IR) spectra (Figure 5a). In comparison with the GO's typical vibrations for C=O ( $1733\text{ cm}^{-1}$ ), aromatic C=C ( $1618\text{ cm}^{-1}$ ), carbonyl (or carboxyl) C-O ( $1411\text{ cm}^{-1}$ ), epoxy C-O ( $1226\text{ cm}^{-1}$ ), and alkoxy C-O ( $1057\text{--}1149\text{ cm}^{-1}$ ), the cl-GO showed a much lower C=O's intensity. This suggests that the GO's energetic epoxy groups were reacted with the  $\text{Al}^{3+}$  cations through the ring-opening reaction, which decreased the content hence intensity of the epoxy vibration. Moreover, the vibration was slightly "red"-shifted to a lower frequency probably because of the leftover ether-like functional groups that are hard to react with the Al (III) cations.

In powder X-ray diffraction (XRD) data (Fig. 5b), the main diffraction peak of the GO sample appear at 2-theta of  $11.4^\circ$  (a lower d-space), whereas that of cl-GO at  $10.39^\circ$  (a higher d-space). This difference proves that upon cross-linking, the d-space in between the stacked cl-GO flakes was increased by the "sandwiched"  $\text{Al}^{3+}$  cations, from 0.79 nm in GO to 0.85 nm in cl-GO. Thus-increased interlayer spacing is a strong evidence of the intercalating-cross-linking of the Al (III) cations in between the cl-GO flakes.

The epoxide ring-opening of GO-polymerization is suggested by X-ray photoelectron spectroscopy (XPS) data of C1s signals of the GO and cl-GO samples (Figure 5c-d). The C-O peak is mainly due to the epoxy/ether groups, and the C=O peak due to the carboxyl and ketone groups. The C-C, C-O and C=O peaks for GO ought to appear at 284.6eV, 286.8eV, and 288.56eV, respectively. The C-C, C-O, and C=O peaks for cl-GO, however, were instead recorded at 284.6eV, 287.0eV, and 288.6eV, respectively. On a much lower intensity, the C-O signal (epoxy/ether peak) of cl-GO significantly shifted upward by 0.2 eV (i.e. more stable or less energetic), which is in line with the epoxide-ring-opening reaction with Al (III) cations.

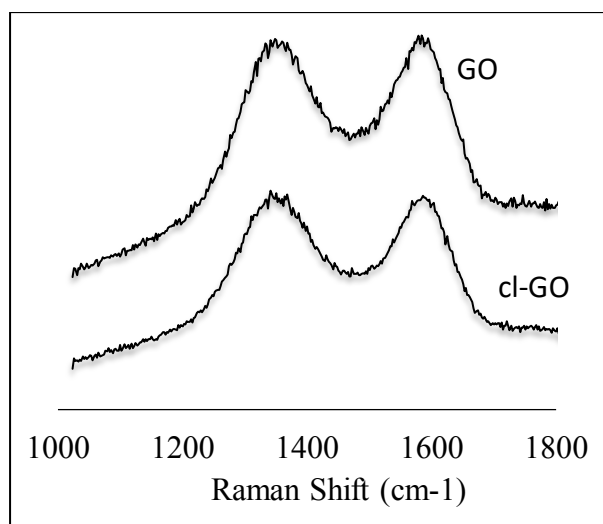




**Figure 5.** Characterizations of GO and cl-GO. a) FT-IR spectra; b) XRD patterns; c) Deconvoluted XPS spectra of GO; d) cl-GO.

The micro-Raman spectra of GO exhibit two broad peaks at  $1593\text{ cm}^{-1}$  and  $1355\text{ cm}^{-1}$  corresponding to the G and D bands, respectively (Figure 6). The G peak is associated with the first order  $E_{2G}$  mode while the D peak is associated with disordered structure of graphite<sup>41,42</sup>. It has been consistently reported that removing oxygenated functional groups from GO can result in an increased D/G signal-intensity ratio, because of the defects from the reduction<sup>43</sup>. Further, the increased D/G signal-intensity ratio has been consistently observed upon reduction of GO,

which could be attributed to the enhanced graphitic structure of GO<sup>41–46</sup>. Accordingly, in this study, the cross-linking changed the D/G signal-intensity ratio, from 1.06 in GO to 1.11 in cl-GO. This slight increase in the D/G signal-intensity ratio of cl-GO suggests that the graphitic character was slightly increased in the cl-GO upon the cross-linking process.



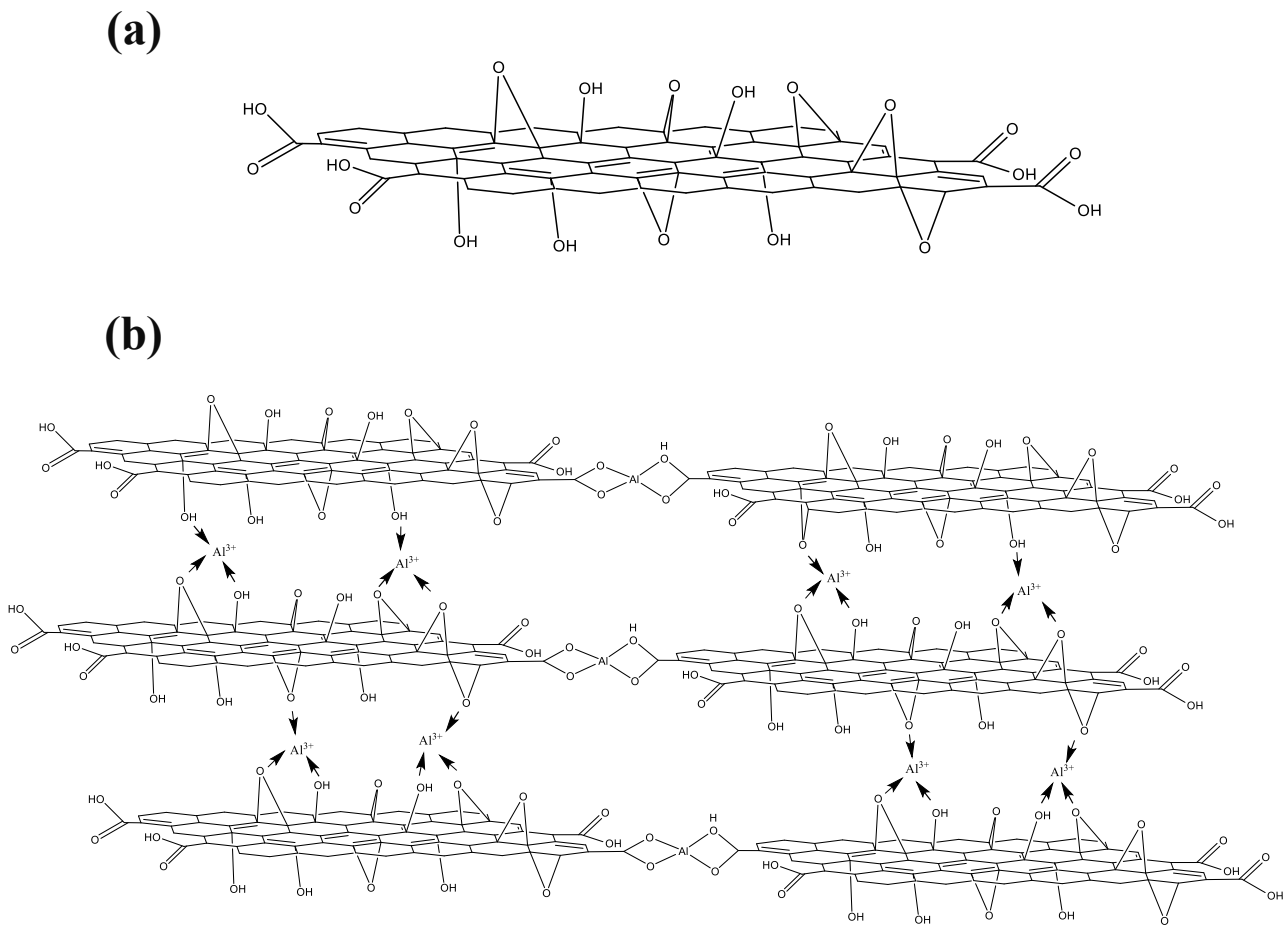
**Figure 6.** Micro-Raman spectra of GO and cl-GO.

In literature,<sup>47</sup> the methylene blue adsorption method is recommended as a simple and effective method to estimate graphitic material's surface area, in which each adsorbed methylene blue's cross-sectional surface area is about 1.35 nm<sup>2</sup>. We, using this method, estimated the surface area of GO and cl-GO to be 645 and 735 m<sup>2</sup>/g, respectively. The extra 90 m<sup>2</sup>/g should be attributed to more methylene blue molecules accessing to the expanded inter-flake space and the Al<sup>3+</sup> cations.

## CONCLUSIONS

We successfully synthesized a new family of nonflammable, water-stable, flexible, lightweight, and mechanically strong polymeric freestanding film of cl-GO, out of the highly flammable, fragile and water-exfoliating GO. All experimental data consistently suggested the

cl-GO to possess such a microstructure as illustrated in Figure 7.



**Figure 7.** Illustration of cl-GO polymer’s microstructure. a) GO; b) cl-GO.

This study confirmed the longstanding and so far not popularly known problem of the high flammability of the as-made GO from the modified Hummer’s method, and developed a facile route to solving the problem that can otherwise dangerously jeopardize the large-scale production and applications of the graphene-related materials. Moreover in materials chemistry, the cross-linking method should be generally applicable to polymerizing many types of layered 2D-materials (e.g. h-BN, MoS<sub>2</sub>, clays, etc.) and even nanocrystals, for meeting new challenge in tailor-making advanced materials at low-cost and high structural precision by design and on-demand.



**ASSOCIATED CONTENT**

**Supporting Information:** The following files are available free of charge. Videos from exposing to open flame with the GO (S1.avi), and cl-GO (S2.avi).

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TOC Graphic

